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## THE PENDING CLAIMS

This listing of claims will replace all prior versions, and listings, of claims in the application.

- 1. (amended) A method of detecting typable loci of a genome, comprising the steps of:
- (a) <u>amplifying genomic DNA with a population of random primers, thereby providing an</u> amplified representative population of genome fragments comprising said typable loci, wherein said population comprises a high complexity representation;
- (b) contacting said genome fragments with a plurality of nucleic acid probes having sequences corresponding to said typable loci under conditions wherein probe-fragment hybrids are formed, wherein said probes are at most 125 nucleotides in length, wherein said nucleic acid probes are immobilized on a substrate; and
  - (c) detecting typable loci of said probe-fragment hybrids.
- 2. (amended) The method of claim 1, wherein said population of representative genome fragments comprises sequences identical to at least 90% [5%] of the genome.
- 3. (original) The method of claim 1, wherein said providing in step (a) comprises representationally amplifying a native genome.
- 4. (original) The method of claim 3, wherein said representationally amplifying comprises using a polymerase of low processivity.
- 5. (original) The method of claim 3, wherein said low processivity is less than 100 bases per polymerization event.
- 6. (original) The method of claim 3, wherein said representationally amplifying comprises a single step reaction yielding a high complexity representation.

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- 7. (original) The method of claim 3, wherein at most  $1 \times 10^6$  copies of said native genome are used as a template for amplification.
  - 8. (canceled)
- 9. (original) The method of claim 8, wherein said substrate is selected from the group consisting of a particle, bead, surface, slide, and microchip.
- 10. (original) The method of claim 1, wherein at least 100 typable loci are simultaneously detected.
  - 11. (original) The method of claim 1, wherein said genome is a human genome.
- 12. (original) The method of claim 1, wherein step (b) comprises contacting said genome fragments with a multiplexed array of nucleic acid probes.
- 13. (amended) The method of claim 12 [1], further comprising contacting said array of nucleic acid probes with chaperone probes.
- 14. (original) The method of claim 1, wherein said probes comprise nucleic acid probes that are at least 20 nucleotides in length.
- 15. (original) The method of claim 1, further comprising producing a report identifying said typable loci that are detected.
  - 16. (canceled)
- 17. (original) The method of claim 1, wherein step (c) comprises directly detecting said typable loci of said fragments that hybridize to said probes.

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- 18. (amended) A method of detecting typable loci of a genome, comprising the steps of:
- (a) <u>amplifying genomic DNA with a population of random primers, thereby providing an</u> amplified representative population of genome fragments comprising said typable loci;
- (b) contacting said genome fragments with a plurality of nucleic acid probes having sequences corresponding to said typable loci under conditions wherein probe-fragment hybrids are formed, wherein said nucleic acid probes are immobilized on a substrate; and
  - (c) directly detecting typable loci of said probe-fragment hybrids
- 19. (amended) The method of claim 18, wherein at most 1000 copies of said [native] genome are amplified.
- 20. (amended) The method of claim 18, wherein said population of representative genome fragments comprises sequences identical to at least 90% [60%] of the genome.
- 21. (original) The method of claim 18, wherein said plurality of nucleic acid probes has sequences for typable loci linked to at least 5% of the expressed sequences of said genome.
- 22. (original) The method of claim 18, wherein said providing in step (a) comprises representationally amplifying a native genome.
- 23. (original) The method of claim 22, wherein said representationally amplifying comprises using a polymerase of low processivity.
- 24. (original) The method of claim 22, wherein said low processivity is less than 100 bases per polymerization event.
- 25. (original) The method of claim 22, wherein said representationally amplifying comprises a single step reaction yielding a high complexity representation.

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- 26. (original) The method of claim 22, wherein at most  $1 \times 106$  copies of said native genome are used as a template for amplification.
  - 27. (canceled)
- 28. (original) The method of claim 18, wherein said substrate is selected from the group consisting of a particle, bead, surface, slide, and microchip.
- 29. (original) The method of claim 18, wherein at least 100 typable loci are simultaneously detected.
  - 30. (original) The method of claim 18, wherein said genome is a human genome.
- 31. (original) The method of claim 18, wherein step (b) comprises contacting said genome fragments with a multiplexed array of nucleic acid probes.
- 32. (original) The method of claim 31, further comprising contacting said array of nucleic acid probes with chaperone probes.
- 33. (amended) The method of claim 18, wherein said probes comprise nucleic acid probes that are at least 20 nucleotides in length.
- 34. (amended) The method of claim 18 [2], further comprising producing a report identifying said typable loci that are detected.
  - 35. (canceled)

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- 36. (original) The method of claim 18, wherein step (c) comprises directly detecting said typable loci of said fragments that hybridize to said probes.
  - 37. (amended) A method of detecting typable loci of a genome, comprising the steps of:
- (a) amplifying genomic DNA with a population of random primers, thereby providing an amplified representative population of genome fragments comprising said typable loci, wherein said population of amplified genome fragments comprises a high complexity representation;
- (b) contacting said genome fragments with a plurality of immobilized nucleic acid probes having sequences corresponding to said typable loci under conditions wherein immobilized probe-fragment hybrids are formed;
  - (c) modifying said immobilized probe-fragment hybrids; and
- (d) detecting a probe or fragment modified in step (c), thereby detecting said typable loci of said genome.
- 38. (original) The method of claim 37, wherein said plurality of nucleic acid probes has sequences for typable loci linked to at least 10% of the expressed sequences of said genome.
- 39. (original) The method of claim 37, wherein said providing in step (a) comprises representationally amplifying a native genome.
- 40. (original) The method of claim 39, wherein said representationally amplifying comprises using a polymerase of low processivity.
- 41. (original) The method of claim 39, wherein said low processivity is less than 100 bases per polymerization event.
- 42. (original) The method of claim 39, wherein said representationally amplifying comprises a single step reaction yielding a high complexity representation.

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- 43. (original) The method of claim 39, wherein at most  $1 \times 10^6$  copies of said native genome are used as a template for amplification.
- 44. (original) The method of claim 37, wherein said nucleic acid probes are immobilized on a substrate.
- 45. (original) The method of claim 44, wherein said substrate is selected from the group consisting of a particle, bead, surface, slide, and microchip.
- 46. (original) The method of claim 37, wherein at least 100 typable loci are simultaneously detected.
  - 47. (original) The method of claim 37, wherein said genome is a human genome.
- 48. (original) The method of claim 37, wherein step (b) comprises contacting said genome fragments with a multiplexed array of nucleic acid probes.
- 49. (original) The method of claim 48, further comprising contacting said array of nucleic acid probes with chaperone probes.
- 50. (original) The method of claim 37, wherein said probes comprises nucleic acid probes are at least 20 nucleotides in length.
- 51. (original) The method of claim 37, further comprising producing a report identifying said typable loci that are detected.
  - 52. (canceled)

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- 53 (original) The method of claim 37, wherein step (c) comprises a primer extension assay.
- 54. (original) The method of claim 53, wherein said primer extension assay is selected from the group consisting of allele specific primer extension (ASPE), single base extension (SBE) and pyrosequencing.

55-63 (canceled)

- 64. (amended) A method for detecting typable loci of a genome, comprising the steps of
- (a) in vitro transcribing a population of amplified genome fragments, thereby obtaining genomic RNA fragments, wherein said population of amplified genome fragments is produced by amplification with a plurality of random primers, wherein said population of amplified genome fragments comprises a high complexity representation;
- (b) hybridizing said genomic RNA fragments with a plurality of nucleic acid probes having sequences corresponding to said typable loci, thereby forming a plurality of RNA fragment-probe hybrids; and
  - (c) detecting typable loci of said RNA fragment-probe hybrids.
  - 65. (canceled)
- 66. (original) The method of claim 64, wherein step (c) comprises modifying said genomic RNA fragment-probe hybrids with reverse transcriptase.
- 67. (original) The method of claim 66, wherein said modifying comprises replicating said genomic RNA fragments hybridized in said genomic RNA fragment-probe hybrids with a plurality of different locus-specific primers, thereby producing a locus-specific, amplified representative population of genome fragments.

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- 68. (original) The method of claim 67, wherein step (a) comprises in vitro transcribing said population of amplified genome fragments using random primers comprising a 3' sequence region that is random and another sequence region having a constant sequence, thereby obtaining genomic RNA fragments labeled with said constant sequence.
- 69. (original) The method of claim 68, wherein said locus-specific primers comprise a 3' sequence region that is locus-specific and a another sequence region having a second constant sequence, thereby obtaining genomic RNA fragments labeled with said first constant region and said second constant region.
- 70. (original) The method of claim 69, further comprising a step of replicating the genomic RNA fragments with complementary primers to the first constant region and second constant region.
- 71. (original) The method of claim 66, wherein said modifying said genomic RNA fragment-probe hybrids with reverse transcriptase occurs under conditions wherein DNA-dependent DNA synthesis is inhibited.
- 72. (original) The method of claim 64, further comprising a step of isolating said genomic RNA fragments.

## 73-77 (canceled)

- 78. (new) The method of claim 1, wherein said genomic DNA is amplified under isothermal conditions using a polymerase having strand displacing activity.
- 79. (new) The method of claim 18, wherein said genomic DNA is amplified under isothermal conditions using a polymerase having strand displacing activity.

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80. (new) The method of claim 37, wherein said genomic DNA is amplified under isothermal conditions using a polymerase having strand displacing activity.